

STIC Search Report

STIC Database Tracking Number: 195114

TO: Ben Sackey

Location: REM 5B31

Art Unit : 1626 July 12, 2006

Case Serial Number: 10/790647

From: Kathleen Fuller Location: EIC 1700

REMSEN 4B28

Phone: 571/272-2505

Kathleen.Fuller@uspto.gov

Search Notes

506 structures from claim structure –only 3 CA references all to the applicants.	
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Scientific and Technical Information Center

SEARCH REQUEST FORM

	- (
Requester's Full Name: 3FN SACE Art Unit: 626 Phone Number: 2-104 Location (Bldg/Room#): (14 5 5 3 Mailbox #): 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Examiner #: 13489 Date: 2/10/03
To ensure an efficient and quality search, please attach a copy of the	e cover sheet, claims, and abstract or fill out the following:
Title of Invention: Novel heterageic (Inventors (please provide full names): Lahray et	ands them prep than containing them
Earliest Priority Date: 7/26/61	·
Search Topic: Please provide a detailed statement of the search topic, and describe as elected species or structures, keywords, synonyms, acronyms, and regis. Define any terms that may have a special meaning. Give examples or r	SCIENTIFIC REFERENCE BR specifically as possible the sequel machinold searthed. Include the stry numbers, and combine with the concept or utility of the invention. relevant citations, authors, etc., if helpen. \hat\(\)
For Sequence Searches Only Please include all pertinent informatio appropriate serial number.	on (parent, child, divisional, or issued patent numbers) along with the Pat. & T.M. Office

R2 R, (CH2) - W-A. P5 P6 7 288

R'-R' are H, halo, OH, S, NH2, CN etc.

CONH3, SONH3, SC2NHMe etc.

R'ad R3 my born 5-6 membered ving

n is 1-8

N is O, S, NR'i rehave R'i is H, C.-2 alkyl a and,

N is O, S, NR'i rehave R'i is H, C.-2 alkyl a and,

I is armate but a wormagehic grap

I is armate but a wormagehic grap

I'm i R'b are H, OH, alkyl, hado

1 is 0, S , is 0, S

; hend's

SACKEY 10/790647 07/12/2006 Page 1

=> FILE REG

FILE 'REGISTRY' ENTERED AT 14:38:45 ON 12 JUL 2006
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 11 JUL 2006 HIGHEST RN 892124-43-5 DICTIONARY FILE UPDATES: 11 JUL 2006 HIGHEST RN 892124-43-5

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TSCA INFORMATION NOW CURRENT THROUGH January 6, 2006

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

=> FILE HCAPLU

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FILE COVERS 1907 - 12 Jul 2006 VOL 145 ISS 3 FILE LAST UPDATED: 11 Jul 2006 (20060711/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> D QUE L8

L1 1 SEA FILE=HCAPLUS ABB=ON US2004-790647/AP

L3 STR

506 structures from quest

REP G1 = (1-10) CH2 VAR G2=O/S/N VAR G3=O/S VAR G4=0/N/S/16 NODE ATTRIBUTES: NSPEC IS R AT 16 DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

N@16

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 16

STEREO ATTRIBUTES: NONE

=> D L8 BIB ABS IND FHITSTR

SSS FUL L3

BB=ON L5

BB=ON L1 AND L7

Applicant with Many Compount

Opplicant with Many Compount

Opplicant with Many Compount

Opplicant with Many Compount

Opplicant with Many Compount 506 SEA FILE=REGISTRY SSS FUL L3 L5

L7 3 SEA FILE=HCAPLUS ABB=ON L5 1 SEA FILE=HCAPLUS ABB=ON L1 AND L7 L8

L8 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 2003:1007858 HCAPLUS

DN 140:59512

TI Preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity

Lohray, Braj Bhushan; Lohray, Vidya Bhushan; Barot, Vijay Kumar Gajubhai; IN Raval, Saurin Khimshanker; Raval, Preeti Saurin; Basu, Sujay

PA Cadilla Healthcare Limited, India

SO U.S. Pat. Appl. Publ., 116 pp., Cont.-in-part of U.S. Pat. Appl. 2003 199,498.

CODEN: USXXCO

 \mathbf{DT} Patent

LA English

FAN. CNT 2

1.171	. CIVI 2						
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
ΡI	US 2003236254	A1	20031225	US 2002-200107	20020719		
	US 7041837	B2	20060509				
	US 2003199498	A1	20031023	US 2001-928242	20010810		
	US 6987123	B2	20060117				

US 2004186099 A1 20040923 US 2004-790647 20040301 <-PRAI IN 2001-MU711 A 20010726
US 2001-928242 A2 20010810
OS MARPAT 140:59512
GI

$$\begin{array}{c|c}
R^{2} & R^{6} & XR^{7} \\
N (CH_{2}) & NWAr & ZR^{8}
\end{array}$$

$$\begin{array}{c|c}
R^{5} & R^{4}
\end{array}$$

AB Title compds. [I; R1-R4 = H, haloalkyl, NO2, cyano, CHO, (substituted)
 alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, aralkyl,
 heterocyclyl, heteroaryl, etc.; W = O, S, NR9; R9 = H, alkyl, aryl; Ar =
 (substituted) aryl, heteroaryl; R5, R6 = H, OH, alkyl, etc.; R5R6 = bond;
 X = O, S; R7 = H, perfluoroalkyl, (substituted) alkyl, cycloalkyl, aryl,
 aralkyl, heteroaryl, heteroaralkyl, heterocyclyl, acyl, etc.; R8 = H,
 (substituted) alkyl, aryl, aralkyl, heteroaryl, heteroaralkyl,
 heterocyclyl, etc.; Y = O, S; Z = O, S, NR10; R10 = H, (substituted)
 alkyl, aryl, aralkyl, hydroxyalkyl, aminoalkyl, heteroaryl, etc.; R8R10 =
 atoms to form a (substituted) 5-6 membered ring; n = 2], were prepared
 Thus, Me 2-ethoxy-3-[6-[2-[2-(4-methoxyphenyl)-5-methylpyrrol-1 yl]ethoxy]naphthalen-2-yl]propanoate (preparation given) at 3 mg/kg day orally
 in mice reduced triglycerides by 26%. I may be useful in the treatment of
 obesity, hyperlipidemia, hypercholesteremia, syndrome X and diabetes.
 Pharmaceutical composition comprising the compound I is claimed.

IC ICM A61K031-541

ICS A61K031-5377; A61K031-496; A61K031-454; A61K031-4439; A61K031-4025; C07D417-02; C07D413-02; C07D043-02

INCL 514227800; X51-425.401; X51-423.55; X51-432.6; X51-440.8; X51-442.2; X54-4 6.0; X54-414.1; X54-437.2; X54-420.8

CC 27-10 (Heterocyclic Compounds (One Hetero Atom))
 Section cross-reference(s): 1, 63

I

ST pyrrolylethoxyphenylethoxypropanoate prepn hypolipemic anticholesteremic; obesity hyperlipidemia hypercholesteremia diabetes treatment pyrrolylethoxyphenylethoxypropanoate prepn

IT Antiarteriosclerotics

(antiatherosclerotics; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT Diabetes mellitus

(complications treatment; preparation of pyrrolylethoxyphenylethoxypropanoat es having hypolipidemic and hypocholesteremic activity)

IT Artery, disease

(coronary, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT Mental and behavioral disorders

(dementia, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT Kidney, disease

(diabetic nephropathy, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

SACKEY 10/790647 07/12/2006 Page 4 IT Eye, disease (diabetic retinopathy, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Blood vessel, disease (endothelium, treatment; preparation of pyrrolylethoxyphenylethoxypropanoate s having hypolipidemic and hypocholesteremic activity) IT Kidney, disease (failure, chronic, treatment; preparation of pyrrolylethoxyphenylethoxypropa noates having hypolipidemic and hypocholesteremic activity) IT Inflammation Kidney, disease (glomerulonephritis, treatment; preparation of pyrrolylethoxyphenylethoxypro panoates having hypolipidemic and hypocholesteremic activity) IT Kidney, disease (glomerulosclerosis, treatment; preparation of pyrrolylethoxyphenylethoxypro panoates having hypolipidemic and hypocholesteremic activity) IT High-density lipoproteins RL: BSU (Biological study, unclassified); BIOL (Biological study) (increasers; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Intestine, disease (inflammatory, treatment; preparation of pyrrolylethoxyphenylethoxypropanoat es having hypolipidemic and hypocholesteremic activity) IT Metabolic disorders (metabolic syndrome X, treatment of; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT. Albumins, biological studies RL: BSU (Biological study, unclassified); BIOL (Biological study) (microalbuminuria, treatment; preparation of pyrrolylethoxyphenylethoxypropa noates having hypolipidemic and hypocholesteremic activity) IT Muscular dystrophy (myotonic, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Kidney, disease (nephrosclerosis, hypertensive nephroslcerosis treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Kidney, disease (nephrotic syndrome, treatment; preparation of pyrrolylethoxyphenylethoxypro panoates having hypolipidemic and hypocholesteremic activity) IT Diabetes mellitus (non-insulin-dependent, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Inflammation Pancreas, disease (pancreatitis, treatment; preparation of pyrrolylethoxyphenylethoxypropanoat es having hypolipidemic and hypocholesteremic activity) ΙŢ Ovary, disease (polycystic, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Anti-inflammatory agents

Antiarteriosclerotics
Anticholesteremic agents
Antidiabetic agents
Antihypertensives
Antiobesity agents
Antitumor agents

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Cardiovascular agents
     Cognition enhancers
     Human
     Hypolipemic agents
        (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic
        and hypocholesteremic activity)
IT
     Low-density lipoproteins
     Very-low-density lipoproteins
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (reducers; preparation of pyrrolylethoxyphenylethoxypropanoates having
        hypolipidemic and hypocholesteremic activity)
IT
     Fatty acids, biological studies
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (reducing free fatty acids; preparation of pyrrolylethoxyphenylethoxypropano
        ates having hypolipidemic and hypocholesteremic activity)
IT
     Glycerides, biological studies
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (reducing plasma triglycerides; preparation of pyrrolylethoxyphenylethoxypro
        panoates having hypolipidemic and hypocholesteremic activity)
IT
     Osteoporosis
        (treatment with antiosteoporotics; preparation of
        pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and
        hypocholesteremic activity)
IT
     Arteriosclerosis
     Atherosclerosis
     Cardiovascular system, disease
     Hypercholesterolemia
     Hyperglycemia
     Hypertension
     Kidney, disease
     Neoplasm
     Obesity
     Psoriasis
     Xanthomatosis
        (treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having
        hypolipidemic and hypocholesteremic activity)
IT
     Dyslipidemia
     Hyperlipidemia
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having
        hypolipidemic and hypocholesteremic activity)
IT
     Endothelium
        (vascular, disease, treatment; preparation of pyrrolylethoxyphenylethoxyprop
        anoates having hypolipidemic and hypocholesteremic activity)
TΤ
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IT

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     494850-04-3P 494850-05-4P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (claimed compound; preparation of pyrrolylethoxyphenylethoxypropanoates
having
       hypolipidemic and hypocholesteremic activity)
     50-78-2, Aspirin
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (coadministration; preparation of pyrrolylethoxyphenylethoxypropanoates
       having hypolipidemic and hypocholesteremic activity)
     50-99-7, D-Glucose, biological studies
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (impaired glucose tolerance treatment; preparation of
       pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and
       hypocholesteremic activity)
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9028-35-7, HMG-CoA reductase
                                                            11128-99-7,
IT
     83-46-5, β-Sitosterol
     Angiotensin II 74315-95-0, \alpha-Glycosidase
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (inhibitors, coadministration; preparation of pyrrolylethoxyphenylethoxyprop
        anoates having hypolipidemic and hypocholesteremic activity)
IT
     169494-85-3, Leptin
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (leptin resistance treatment; preparation of pyrrolylethoxyphenylethoxypropa
        noates having hypolipidemic and hypocholesteremic activity)
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RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic
   and hypocholesteremic activity)
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IT

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RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic
   and hypocholesteremic activity)
                               106-93-4, 1,2-Dibromoethane
                                                              107-21-1,
105-36-2, Ethyl bromoacetate
                             123-08-0, 4-Hydroxybenzaldehyde
Ethylene glycol, reactions
                                                                141-43-5,
                         1003-29-8, 2-Formylpyrrole
Ethanolamine, reactions
                                                       4437-46-1,
1-Phenylhexane-1,4-dione 13676-06-7, Triethyl 2-ethoxyphosphonoacetate
53391-61-0, 2-Methylthiopyrrole
                                  197299-16-4, Ethyl 3-(4-hydroxyphenyl)-2-
                   222555-06-8, Ethyl (S)-3-(4-hydroxyphenyl)-2-
ethoxypropionate
                   267228-43-3, Methyl (S)-3-(4-hydroxyphenyl)-2-
ethoxypropionate
                    494852-05-0
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methoxypropionate
RL: RCT (Reactant); RACT (Reactant or reagent)
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   and hypocholesteremic activity)
6719-02-4P, 1H-Pyrrole-1-ethanol
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                                              494852-04-9P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
   (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic
   and hypocholesteremic activity)
59-67-6, Nicotinic acid, biological studies
                                               9004-10-8, Insulin,
```

biological studies 11041-12-6, Cholestyramine 23288-49-5, Probucol 50925-79-6, Cholestipol

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT 494848-08-7P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(claimed compound; preparation of pyrrolylethoxyphenylethoxypropanoates

having

hypolipidemic and hypocholesteremic activity)

RN 494848-08-7 HCAPLUS

CN Benzenepropanamide, α-ethoxy-N-[(1S)-2-hydroxy-1-phenylethyl]-4-[2-(2-methyl-5-phenyl-1H-pyrrol-1-yl)ethoxy]-, (αR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

=> D QUE

L1 L3 1 SEA FILE=HCAPLUS ABB=ON US2004-790647/AP STR

N @16

REP G1=(1-10) CH2

VAR G4=0/N/S/16 NODE ATTRIBUTES: NSPEC IS R

AT 16 DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 16

STEREO ATTRIBUTES: NONE

506 SEA FILE=REGISTRY SSS FUL L3 L5 L7 3 SEA FILE=HCAPLUS ABB=ON L5

1 SEA FILE=HCAPLUS ABB=ON L1 AND L7 $^{\text{L8}}$ 2 SEA FILE=HCAPLUS ABB=ON L7 NOT L8 L9

=> SEL HIT RN L9 1-2 E1 THROUGH E360 ASSIGNED Remaining 2 references 360 structures so printed only 1 structure for each

=> D L9 BIB ABS IND FHITSTR 1-2

ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN L9

2003:97297 HCAPLUS AN

DN 138:153432

ΤI Preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity.

Lohray, Braj Bhushan; Lohray, Vidya Bhushan; Barot, Vijay Kumar; Raval / applicante Saurin Khimshankar; Raval, Preeti Saurin; Basu, Sujay IN Saurin Khimshankar; Raval, Preeti Saurin; Basu, Sujay Cadila Healthcare Limited, India

PA

SO PCT Int. Appl., 116 pp.

CODEN: PIXXD2

DT Patent

LA English FAN CNT 2

PATENT NO.						KIND DATE			APPLICATION NO.						DATE				
ΡI	WO 2003009841			A1 · 20030206								20020725							
		W:										BG,							
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	
			PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,	
			UA,	UG,	US,	UΖ,	VN,	YU,	ZA,	ZM,	ZW								
		RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	ŪĠ,	ZM,	ZW,	AT,	BE,	BG,	
			CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	
			PT,	SE,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	
			ΝE,	SN,	TD,	TG													
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	EΡ	1414	439			A1 20040506				EP 2002-751609						20020725			
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
			ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	SK			
	CN	1558	758			Α		2004	1229		ČN 2002-818983					20020725			
	BR	2002	0116	65		Α		20050111		:	BR 2002-1		-11665			20020725			
	JP	2005	5033	67		T2		2005	0203		JP 2003-53		515234			20020725			
	ИО	2004	0003	01		Α		20040324		1	NO 2004		2004-301			20	0040	123	
	ZA	2004	0005	63		Α		2004	0041029 ZA 2004-56				563			20	040	126	
PRAI	IN	2001	-MU7	11		Α		2001	0726										

WO 2002-IN155 MARPAT 138:153432 W 20020725

OS GT

$$\begin{array}{c|c}
R1 & R6 & XR7 & Y \\
N (CH2) nWAr & ZR8 \\
R4 & R5
\end{array}$$

AB Title compds. [I; R1, R4 = H, haloalkyl, NO2, cyano, CHO, (substituted)
alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, aralkyl,
heterocyclyl, heteroaryl, etc.; W = O, S, NR9; R9 = H, alkyl, aryl; Ar =
 (substituted) aryl, heteroaryl; R5, R6 = H; R5R6 = bond; R7 = H,
 perfluoroalkyl, (substituted) alkyl, cycloalkyl, aryl, aralkyl,
heteroaryl, heteroaralkyl, heterocyclyl, acyl, etc.; R8 = H, (substituted)
alkyl, aryl, aralkyl, heteroaryl, heteroaralkyl, heterocyclyl, etc.; Y =
 O, S; Z = O, S, NR10; R10 = H, (substituted) alkyl, aryl, aralkyl,
 hydroxyalkyl, aminoalkyl, heteroaryl, etc.; R8R10 = atoms to form a
 (substituted) 5-6 membered ring; n = 2], were prepared Thus, Me
 2-ethoxy-3-[6-[2-[2-(4-methoxyphenyl)-5-methylpyrrol-1 yl]ethoxy]naphthalen-2-yl]propanoate (preparation given) at 3 mg/kg day orally
 in mice reduced triglycerides by 26%. I may be useful in the treatment of
 obesity, hyperlipidemia, hypercholesteremia, syndrome X and diabetes.

IC ICM A61K031-40 ICS C07D207-325; C07D207-333; C07D407-04; C07D409-04; C07D401-04; A61K031-4025; A61P003-06

CC 27-10 (Heterocyclic Compounds (One Hetero Atom))
 Section cross-reference(s): 1

Ι

pyrrolylethoxyphenylethoxypropanoate prepn hypolipidemic hypocholesteremic; obesity hyperlipidemia hypercholesteremia diabetes treatment pyrrolylethoxyphenylethoxypropanoate prepn

IT Diabetes mellitus

(complications treatment; preparation of pyrrolylethoxyphenylethoxypropanoat es having hypolipidemic and hypocholesteremic activity)

IT Mental and behavioral disorders

(dementia, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT Kidney, disease

(diabetic nephropathy, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT Eye, disease

(diabetic retinopathy, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT Blood vessel, disease

(endothelium, treatment; preparation of pyrrolylethoxyphenylethoxypropanoate s having hypolipidemic and hypocholesteremic activity)

IT Kidney, disease

(failure, chronic, treatment; preparation of pyrrolylethoxyphenylethoxypropa noates having hypolipidemic and hypocholesteremic activity)

IT Inflammation

Kidney, disease

(glomerulonephritis, treatment; preparation of pyrrolylethoxyphenylethoxypro

panoates having hypolipidemic and hypocholesteremic activity) IT Kidney, disease (glomerulosclerosis, treatment; preparation of pyrrolylethoxyphenylethoxypro panoates having hypolipidemic and hypocholesteremic activity) High-density lipoproteins IT RL: BSU (Biological study, unclassified); BIOL (Biological study) (increasers; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Intestine, disease (inflammatory, treatment; preparation of pyrrolylethoxyphenylethoxypropanoat es having hypolipidemic and hypocholesteremic activity) IT Albumins, biological studies RL: BSU (Biological study, unclassified); BIOL (Biological study) (microalbuminuria, treatment; preparation of pyrrolylethoxyphenylethoxypropa noates having hypolipidemic and hypocholesteremic activity) IT Muscular dystrophy (myotonic, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Kidney, disease (nephrosclerosis, hypertensive nephroslcerosis treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Kidney, disease (nephrotic syndrome, treatment; preparation of pyrrolylethoxyphenylethoxypro panoates having hypolipidemic and hypocholesteremic activity) IT Diabetes mellitus (non-insulin-dependent, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Inflammation Pancreas, disease (pancreatitis, treatment; preparation of pyrrolylethoxyphenylethoxypropanoat es having hypolipidemic and hypocholesteremic activity) IT Ovary, disease (polycystic, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) TT Antiarteriosclerotics Anticholesteremic agents Antidiabetic agents Antiobesity agents Antitumor agents Cognition enhancers Human Hypolipemic agents (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Low-density lipoproteins Very-low-density lipoproteins RL: BSU (Biological study, unclassified); BIOL (Biological study) (reducers; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Osteoporosis (treatment with antiosteoporotics; preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity) IT Arteriosclerosis Cardiovascular system, disease Hypercholesterolemia Hyperglycemia Kidney, disease

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Neoplasm
     Obesity
     Psoriasis
     Xanthomatosis
        (treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having
        hypolipidemic and hypocholesteremic activity)
IT
     Hyperlipidemia
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (treatment; preparation of pyrrolylethoxyphenylethoxypropanoates having
        hypolipidemic and hypocholesteremic activity)
IT
     Endothelium
        (vascular, disease, treatment; preparation of pyrrolylethoxyphenylethoxyprop
        anoates having hypolipidemic and hypocholesteremic activity)
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495399-38-7P 495399-55-8P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic
   and hypocholesteremic activity)
105-36-2, Ethyl bromoacetate 106-93-4, 1,2-Dibromoethane
                                                              107-21-1,
Ethylene glycol, reactions
                             123-08-0, 4-Hydroxybenzaldehyde
                                                                141-43-5,
                         1003-29-8, 2-Formylpyrrole 4437-46-1,
Ethanolamine, reactions
                          13676-06-7, Triethyl 2-ethoxyphosphonoacetate
1-Phenylhexane-1,4-dione
53391-61-0, 2-Methylthiopyrrole
                                 197299-16-4, Ethyl 3-(4-hydroxyphenyl)-2-
ethoxypropionate 222555-06-8, Ethyl (S)-3-(4-hydroxyphenyl)-2-
                   267228-43-3, Methyl (S)-3-(4-hydroxyphenyl)-2-
ethoxypropionate
                    494852-05-0
methoxypropionate
                                  494852-06-1
RL: RCT (Reactant); RACT (Reactant or reagent)
   (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic
   and hypocholesteremic activity)
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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT 59-67-6, Nicotinic acid, biological studies 9004-10-8, Insulin, biological studies 11041-12-6, Cholestyramine 23288-49-5, Probucol 50925-79-6, Cholestipol

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of pyrrolylethoxyphenylethoxypropanoates having hypolipidemic and hypocholesteremic activity)

IT 494848-08-7P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(claimed compound; preparation of pyrrolylethoxyphenylethoxypropanoates.

having

hypolipidemic and hypocholesteremic activity)

RN 494848-08-7 HCAPLUS

CN Benzenepropanamide, α-ethoxy-N-[(1S)-2-hydroxy-1-phenylethyl]-4-[2-(2-methyl-5-phenyl-1H-pyrrol-1-yl)ethoxy]-, (αR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 2001:545659 HCAPLUS

DN 135:137396

TI Preparation of pyrrolylethoxyphenylethoxypropanoates and related compounds for treatment of hyperglycemia, hypertension, cardiovascular disease, and eating disorders.

IN Lohray, Braj Bhushan; Loray, Vidya Bhushan; Barot, Vijay Kumar Gajubhai

PA Cadila Healthcare Ltd., India

SO PCT Int. Appl., 54 pp.

CODEN: PIXXD2

DT Patent

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			HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT
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MARPAT 135:137396

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BG 106932

PRAI IN 2000-MU57

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AB Title compds. [I; R1-R4 = H, halo, perhaloalkyl, OH, SH, amino, NO2, etc.; R2R3 = atoms to form a (substituted) 5-6 membered (heterocyclic) ring; R5, R6 = H, or R5R6 = bond, or R5, R6 = OH, alkyl, alkoxy, halo, acyl, (substituted) aralkyl; X, Y = O, S; R7 = H, perfluoroalkyl, (substituted) alkyl, cycloalkyl, aryl, aralkyl, heteroaryl, heterocyclyl, alkoxyalkyl, aryloxyalkyl, etc.; W = O, S, NR9; Z = O, NR10; R8 = H, (substituted) alkyl, aryl, aralkyl, heteroaryl, heteroaralkyl, heterocyclyl, hydroxyalkyl, etc.; R9 = alkyl, aryl; R10 = H, (substituted) alkyl, aryl, aralkyl, hydroxyalkyl, aminoalkyl, heteroaryl, etc.; Ar = (substituted) (fused) divalent aryl, heteroaryl, heterocyclyl], were prepared as drugs (no data). Thus, Et 3-(4-hydroxyphenyl)-2-ethoxypropanoate, K2CO3, and DMF were stirred at 70-80° for 10 min. followed by addition of 2-(2,5-dimethyl-1H-pyrrol-1-yl)ethyl methanesulfonate (preparation given) followed by stirring for 5 h at 70-80° and standing overnight to give 89% Et 3-[4-[2-(2,5-dimethylpyrrol-1-yl)ethoxy]phenyl]-2ethoxypropanoate.

351426-21-6P

TT

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IC
     C07D207-00
CC
     27-10 (Heterocyclic Compounds (One Hetero Atom))
     Section cross-reference(s): 1
ST
     pyrrolylethoxyphenylethoxypropanoate prepn hyperglycemia hypertension
     cardiovascular disease eating disorder treatment
IT
     Appetite
        (disorder, treatment; preparation of pyrrolylethoxyphenylethoxypropanoates
        and related compds. for treatment of hyperglycemia, hypertension,
        cardiovascular disease, and eating disorders)
IT
     Antidiabetic agents
     Antihypertensives
     Cardiovascular agents
        (preparation of pyrrolylethoxyphenylethoxypropanoates and related compds.
        for treatment of hyperglycemia, hypertension, cardiovascular disease,
        and eating disorders)
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     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (preparation of pyrrolylethoxyphenylethoxypropanoates and related compds.
        for treatment of hyperglycemia, hypertension, cardiovascular disease,
        and eating disorders)
     110-13-4, Hexane-2,5-dione
                                  141-43-5, Ethanolamine, reactions
TΤ
     197299-16-4
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of pyrrolylethoxyphenylethoxypropanoates and related compds.
        for treatment of hyperglycemia, hypertension, cardiovascular disease,
        and eating disorders)
TT
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     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of pyrrolylethoxyphenylethoxypropanoates and related compds.
        for treatment of hyperglycemia, hypertension, cardiovascular disease,
       and eating disorders)
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RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);

BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of pyrrolylethoxyphenylethoxypropanoates and related compds. for treatment of hyperglycemia, hypertension, cardiovascular disease, and eating disorders)

RN 351426-21-6 HCAPLUS

CN Benzenepropanoic acid, $4-[2-(2,5-dimethyl-1H-pyrrol-1-yl)ethoxy]-\alpha-ethoxy-, ethyl ester (9CI) (CA INDEX NAME)$

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